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Journal of Pharmaceutical Research and Development (1998), 3(2), 75-79. Ekpendu, T. O. E etal. Chemical constituents of the extractives of Napoleona imperialis P. Beauv. (Lecythidaceae).

ACTA CRYSTALLOGR SECT B STRUCT CRYSTALLOGR CRYST CHEM, (1980) 36 (7), 1593-1598.). SPIRLET M R etal. STRUCTURE OF ACETYLATED NAPOLEOGENIN.

Phytochemistry (1980), 19(4), 615-22. Kapundu, Mpuza etal. New triterpenoids from Napoleonaea imperialis.

67) 634 AZL 10/17-RC и В36 PART 7 Pp. 1525-1732

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Acta Crystallographica

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The Structure of Acetylated Napoleogenin

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Abstract

Acceylated napoleogenin, $C_{58}H_{84}O_{18}$, crystallizes in sace group $P2_1$ with two molecules in a unit cell of imensions $a=19\cdot127$ (1), $b=10\cdot812$ (1), c=1.879 (1) Å, $\beta=102\cdot540$ (2)°. The crystal structure as solved by direct methods. Full-matrix least-squares refinement with all atoms treated isotropically and sing 3139 reflections gave an R value of $0\cdot108$. The compound possesses a triterpene skeleton with cissed D and E rings in the chair form. A 3,4-0-angelyl-6-deoxy- β -glucopyranosyl group sits at (21) on the E ring. The molecular structure of

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acetylated napoleogenin is formulated by the present analysis as 21β -(2-O-acetyl-3,4-di-O-angelyl-6-deoxy- β -glucopyranosyloxy)-3 β ,16 α ,22 α ,24,28-pentaacetoxy-olean-12-ene. The large distortion of the whole molecular skeleton may result from 1,3 diaxial interaction. The packing is mainly dictated by close van der Waals contacts between the molecules.

Introduction

Within the framework of chemical taxonomy studies, the acetylated form of napoleogenin, a new pentacyclic © 1980 International Union of Crystallography

O(4')

O(5')

0.9187(4)

0.8376(6)

triterpene, was isolated from the seeds of Napoleonaea imperialis, a plant native to Zaïre. An X-ray crystal-structure investigation of the compound was undertaken in order to establish unequivocally its chemical structure and stereochemistry.

Table 1. Atomic positional and isotropic thermal parameters with standard deviations in acetylated napoleogenin

(i) Triterpene nucleus C(1)		x	y	z	$B(\dot{A}^2)$
C(1)	(i) Tritor	nene nucleus			
C(2) 0.5595 (8) 0.063 (2) -0.164 (1) 7.2 (3) C(3) 0.6094 (7) -0.002 (2) -0.214 (1) 6.1 (3) C(4) 0.6195 (7) -0.142 (2) -0.1966 (9) 5.9 (3) C(5) 0.6442 (6) -0.159 (1) -0.1917 (8) 5.1 (2) C(6) 0.6609 (6) -0.290 (1) -0.0571 (8) 4.8 (2) C(7) 0.7072 (7) -0.287 (1) 0.0403 (9) 5.5 (3) C(8) 0.6706 (6) -0.222 (1) 0.1111 (7) 4.4 (2) C(9) 0.6418 (6) -0.094 (1) 0.0712 (7) 4.4 (2) C(9) 0.6418 (6) -0.094 (1) 0.0712 (7) 4.4 (2) C(10) 0.5957 (6) -0.096 (1) -0.0314 (8) 4.8 (2) C(11) 0.6039 (7) -0.027 (2) 0.136 (1) 6.8 (3) C(12) 0.6383 (7) -0.045 (2) 0.2365 (9) 5.7 (3) C(13) 0.6943 (6) -0.124 (1) 0.2691 (8) 4.8 (2) C(14) 0.7283 (6) -0.196 (1) 0.2482 (8) 5.1 (2) C(16) 0.7920 (6) -0.319 (1) 0.2482 (8) 5.1 (2) C(16) 0.7920 (6) -0.319 (1) 0.2482 (8) 5.1 (2) C(16) 0.7920 (6) -0.319 (1) 0.3532 (8) 4.7 (2) C(17) 0.7468 (6) -0.253 (1) 0.4107 (7) 4.4 (2) C(19) 0.7756 (5) -0.022 (1) 0.4064 (7) 4.4 (2) C(20) 0.8003 (6) -0.124 (1) 0.5115 (8) 4.6 (2) C(21) 0.8372 (5) -0.137 (1) 0.5414 (7) 3.8 (2) C(22) 0.7860 (6) -0.242 (1) 0.5156 (7) 4.5 (2) (2) 0.7890 (6) -0.319 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.319 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.124 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.124 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.124 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.124 (1) 0.5115 (8) 4.6 (2) (2) 0.7890 (6) -0.124 (1) 0.5156 (7) 4.5 (2) (2) 0.7890 (6) -0.124 (1) 0.5156 (7) 4.5 (2) (2) 0.7890 (9) 0.132 (2) 0.5577 (9) 5.7 (3) (2) (2) 0.553 (1) 0.092 (2) 0.5577 (9) 5.7 (3) (2) (2) 0.553 (1) 0.017 (1) 0.030 (2) 0.5577 (9) 5.7 (3) (2) 0.599 (9) 0.132 (2) 0.0348 (1) 7.7 (4) (2) (2) 0.553 (1) 0.013 (2) 0.1299 (9) 5.8 (3) (2) 0.059 (2) 0.5577 (9) 5.7 (3) (2) 0.6493 (9) 0.132 (2) 0.0348 (1) 7.7 (4) (2) (2) 0.6493 (9) 0.132 (2) 0.0348 (1) 7.7 (4) (2) (2) 0.6359 (7) 0.020 (2) 0.5577 (9) 5.7 (3) (2) (2) 0.6493 (9) 0.132 (2) 0.0348 (1) 7.7 (4) (2) 0.6493 (9) 0.032 (2) 0.0379 (1) 0.040 (1) 0.6493 (9) 0.056 (2) 0.0311 (1) 0.0565 (5) 0.040 (1) 0.580 (1) 0.0565 (2) 0.040 (1) 0.066 (1) 0.958 (1) 0.0565 (2) 0.040 (1) 0.066 (1) 0.958 (1) 0.0565 (2			0.047(1)	-0.0589 (9)	6.3 (3)
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(ii) Extranuclear (C(23)		` '		` '	
(ii) Extranuclear C(23)		• •			
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O(11) $0.8596(4)$ $-0.138(1)$ $0.6416(5)$ $4.5(1)$		0.7681 (7)		0.6413 (8)	9.0(3)
	O(11)	0.8596 (4)	-0.138(1)	0.6416 (5)	4.5 (1)

		y	٠,	D (A)
(iii) Gluc	opyranosyl mo	oiety		
Č(1')	0.9320 (6)	-0.128(1)	0-6775 (7)	4.3 (1
C(2')	0.9452 (6)	-0.079(1)	0.7747 (8)	4.5 (1
C(3')	1.0269 (5)	-0.072(1)	0-8153 (8)	4-7(
C(4')	1-0603 (6)	-0·192 (1)	0.8042 (8)	4.7 (Our
C(5')	1.0391 (7)	-0.240(1)	0:7026 (9)	5.60 genin
O(1')	0.9637 (4)	-0.247(1)	0.6774 (5)	
C(6')	1.0692 (8)	-0.370(2)	0.691 (1)	6.40 B-am
C(7')	1.1814 (8)	-0.212(2)	0.897 (1)	6.40 gluco
C(8')	1.260(1)	-0·187 (2)	0.893 (1)	8-2 More
C(9')	1.301 (1)	-0.156 (3)	0.977 (2)	11-2 grou
C(10')	1.2873 (7)	-0·142 (2)	1.0605 (9)	9·1 (at C
C(11')	1.279 (1)	-0 ⋅192 (3)	0.798 (2)	11.7(4
C(7")	1.0759 (8)	0.047 (2)	0.948 (1)	6.50 Blocat
C(8")	1.078 (1)	0.061 (2)	1.053 (1)	8-8 meth
C(9")	1.0416 (7)	0.015 (2)	1.0960 (9)	8.0 dedu
C(10")	0.9722 (7)	-0.058 (2)	1-0686 (9)	9-10 lated
C(11")	1.142 (1)	0.149 (2)	1-093 (1)	X X D
C(12')	0.8652 (7)	0.064 (2)	0-817 (1)	6.5 (Egluco
C(13')	0.840(1)	0.196 (2)	0-807 (1)	8.8 (olean
O(2')	1-1383 (4)	-0·172 (1)	0-8199 (5)	5.6(Fin F
O(3')	1.1558 (8)	-0·258 (2)	0.956(1)	11.0(in Fi
O(2")	1.0318 (4)	-0.051 (1)	0.9124 (6)	5.6 (PLU
O(3")	1-1033 (8)	0.116(2)	0.907(1)	10.0

Table 1 (cont.)

Experimental

Colorless needle-shaped crystals, elongated along the axis, were grown from methanol.

0.044(1)

-0.014 (1)

0.7743(5)

0.8567(9)

deposi tary P

throug lograp

The space group was unequivocally established by Weissenberg and precession photographs.

Cell parameters and intensities were measured with four-circle Hilger diffractometer using Cu $K\alpha$ radiatio ($\lambda = 1.5418$ Å, $\omega - 2\theta$ scan). Integrated intensities were recorded for 3353 independent reflections. They were corrected for Lorentz and polarization effects but not for absorption (μ for Cu $K\alpha = 0.72$ mm⁻¹).

The structure was solved by direct methods using MULTAN (Main, Lessinger, Woolfson, Germain) Declercq, 1977). A large number of cycles of structure factor and Fourier calculations were necessary to fig all the atoms. The oxygen atoms were identified w the help of chemical and structural consideration Refinement of atomic coordinates and individual isotropic thermal parameters for C and O atoms w carried out by a full-matrix least-squares method (program SFLS, Prewitt, 1967) on 3139 reflections which $I > 2\sigma(I)$. In the last cycles of refinement, atoms were included in calculated positions assumi C-H bond lengths of 1.087 A. All were given same isotropic temperature factor $B = 4.0 A^{23}$ Cruickshank (1961) type of weighting scheme used. The final value of the conventional discrepand factor R is 0.108. We could not carry out further refinement with anisotropic thermal parameters: large number of variables would have involved

checomputing time and expense. Our final positional in the differential parameters are listed in Table 1.*

Discussion

(7), (8), (8), (8), (9), (5)

2) (9)

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4-3 (2)

4·5 (2) 4·7 (2)

10-0 (3)

5-4 (2) 8-9 (3)

ted along the b

established by

easured with u Ka radiation intensities were ns. They were effects but not methods using ų, Germain & les of structure cessary to find identified with considerations and individual 1. O atoms was juares meth^{od} 9 reflections for refinement, H itions assuming were given the $= 4.0 \text{ Å}^2. \text{ }^{\text{A}}$ g scheme was nal discrepanci rry out furths parameters: the e involved too

Our structural results indicate that acetylated napoleomin is a pentacyclic oleanene-type triterpene of the myrin series with a 3,4-di-o-angelyl-6-deoxy-βpucopyranosyl group located on the E ring at C(21). Moreover, in the triterpene nucleus, a geminal dimethyl group is situated at C(20) and four other methyl groups at C(4), C(8), C(10) and C(14). Acetyl groups are located at C(3), C(16) and C(22) while two acetoxymethyl groups are found at C(4) and C(17). We deduced that acetylated napoleogenin should be formulated as 21β -(2-O-acetyl-3,4-di-O-angelyl-6-deoxy- β glucopyranosyloxy) - 3β , 16α , 22α , 24, 28 - pentaacetoxyokan-12-ene. The numbering of the atoms is illustrated n-Fig. 1. The molecular conformation is shown in Fig. 2, a stereodrawing produced by the program PLUTO (Crystallographic Data Files, 1979). Bond lengths and angles are listed in Tables 2 and 3.

Lists of structure factors and hydrogen coordinates have been deposited with the British Library Lending Division as Supplementary Publication No. SUP .35161 (19 pp.). Copies may be obtained through The Executive Secretary, International Union of Crystallography, 5 Abbey Square, Chester CH1 2HU, England.

Fig. 1. Numbering of atoms in acetylated napoleogenin.

Table 2. Bond lengths (A) with standard deviations in parentheses

(i) Triterpene nucleus		(ii) Extranuclear	
C(1)-C(2)	1.54 (2)	C(3)-O(1)	1.47 (2)
C(1)-C(10)	1.60 (2)	C(4)—C(23)	1.56 (2)
C(2)-C(3)	1.51 (2)	C(4)-C(24)	1.51 (2)
C(3)-C(4)	1.54(2)	C(8)-C(26)	1.51 (2)
C(4)-C(5)	1.54 (2)	C(10)-C(25)	1.54 (2)
C(5)-C(6)	1.52 (2)	C(14)-C(27)	1.56 (2)
C(5)-C(10)	1.58 (2)	C(16)-O(5)	1.48 (1)
C(6)-C(7)	1.53 (2)	C(17)-C(28)	1.54 (2)
C(7)-C(8)	1.56 (2)	C(20)C(29)	1.54 (2)
C(8)-C(9)	1.56 (2)	C(20)-C(30)	1.52 (2)
C(8)-C(14)	1.59 (2)	C(21)-O(11)	1.46 (1)
C(9)-C(10)	1.59 (2)	C(22)-O(9)	1.45 (2)
C(9)-C(11)	1.51 (2)	C(31)-O(1)	1.40(2)
C(11)-C(12)	1.51(2)	C(31)=O(2)	1.11(2)
C(12)=C(13)	1.37(2)	C(31)-C(32)	1-50 (3)
C(13)-C(14)	1.50(2)	C(33)-O(3)	1.34(2)
C(13)-C(18)	1.53 (2)	C(33)=O(4)	1.27(2)
C(14)-C(15)	1.52 (2)	C(33)-C(34)	1.42 (3)
C(15)-C(16)	1.56 (2)	C(35)-O(5)	1.34 (2)
C(16)-C(17)	1.52 (2)	C(35)=O(6)	1.21 (2)
C(17)-C(18)	1.54 (2)	C(35)-C(36)	1.45 (3)
C(17)-C(22)	1.58 (2)	C(37)—O(7)	1.31(2)
C(18)-C(19)	1.53 (2)	C(37) = O(8)	1.30 (3)
C(19)-C(20)	1.54 (2)	C(37)-C(38)	1.46 (3)
C(20)C(21)	1.56 (2)	C(39)-O(9)	1.40(2)
C(21)-C(22)	1.50(2)	C(39)=O(10)	1.08(2)
0(21) 0(22)	1 50 (2)	C(39)-C(40)	1.55 (3)
		-()	
(iii) Glucopyran	osvl moiety		
C(1')-O(11)	1-38(1)	C(7')-C(8')	1.54 (2)
C(1')-O(1')	1.42 (2)	C(7'')-O(2'')	1.38 (2)
C(1')-C(2')	1·51 (2)	C(7'')=O(3'')	1.16 (2)
C(2')-O(4')	1.42 (1)	C(7")-C(8")	1.56 (3)
C(2')-C(3')	1·55 (2)	C(8')-C(11')	1.53 (3)
C(3')-O(2'')	1.45 (1)	C(8')=C(9')	1.38 (3)
C(3')-C(4')	1.48 (2)	C(8'')-C(11'')	1.56 (3)
C(4')-O(2')	1.48 (1)	C(8")=C(9")	1.15 (3)
C(4')C(5')	1.57(2)	C(9')-C(10')	1.53 (2)
C(5')-O(1')	1.41(1)	C(9'')-C(10'')	1.53 (2)
C(5')C(6')	1.54 (2)	C(12')-O(4')	1.34 (2)
C(7')-O(2')	1.34(2)	C(12')=O(5')	1.21 (2)
C(7')=O(3')	1.20(2)	C(12')-C(13')	1.52 (3)
. , ,	,		

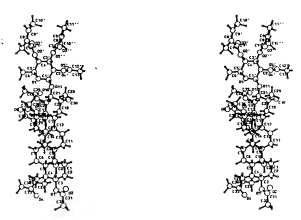


Fig. 2. Stereoscopic drawing of the molecule.

Table 3. Bond angles (°) with standard deviations in parentheses

(i) Triterpene nucleus			
C(2)-C(1)-C(10)	111 (1)	C(12)=C(13)-C(18)	115 (1)
C(1)-C(2)-C(3)	109 (1)	C(14)-C(13)-C(18)	124 (1)
C(2)-C(3)-C(4)	117 (1)	C(8)-C(14)-C(13)	110 (1)
C(2)-C(3)-O(1)	107 (1)	C(8)-C(14)-C(15)	110(1)
	108 (1)	C(8)—C(14)—C(27)	112(1)
C(4)-C(3)-O(1)			
C(3)-C(4)-C(5)	106 (1)	C(13)-C(14)-C(15)	110(1)
C(3)-C(4)-C(23)	109 (1)	C(13)-C(14)-C(27)	106 (1)
C(3)-C(4)-C(24)	108 (1)	C(15)-C(14)-C(27)	109 (1)
C(5)-C(4)-C(23)	107 (1)	C(14)C(15)C(16)	118 (1)
C(5)-C(4)-C(24)	117 (1)	C(15)C(16)C(17)	113 (1)
C(23)-C(4)-C(24)	108 (1)	C(15)-C(16)-O(5)	108-0 (9)
C(4)-C(5)-C(6)	117 (1)	C(17)-C(16)-O(5)	107 (1)
C(4)-C(5)-C(6) C(4)-C(5)-C(10)	116 (1)	C(16)-C(17)-C(18)	114(1)
C(4)-C(3)-C(10)	108 (1)	C(16)-C(17)-C(22)	113 (1)
C(6)-C(5)-C(10)			107 (1)
C(5)-C(6)-C(7)	110 (1)	C(16)—C(17)—C(28)	
C(6)-C(7)-C(8)	114 (1)	C(18)C(17)C(22)	110(1)
C(7)-C(8)-C(9)	109 (1)	C(18)-C(17)-C(28)	105 (1)
C(7)-C(8)-C(14)	110(1)	C(22)-C(17)-C(28)	109 (1)
C(7)-C(8)-C(26)	110(1)	C(13)-C(18)-C(17)	112(1)
C(9)C(8)C(14)	107-3 (9)	C(13)-C(18)-C(19)	112(1)
C(9)-C(8)-C(26)	111 (1)	C(17)-C(18)-C(19)	113(1)
C(14)-C(8)-C(26)	110 (1)	C(18)-C(19)-C(20)	115 (1)
C(8)-C(9)-C(10)	115 (1)	C(19)-C(20)-C(21)	104 (1)
C(8)-C(9)-C(11)	iii (i)	C(19)-C(20)-C(29)	108 (1)
C(10) C(1) C(11)		C(19) C(20) C(20)	112(1)
C(10)-C(9)-C(11)	112 (1)	C(19)-C(20)-C(30)	
C(1)-C(10)-C(5)	108 (1)	C(21)-C(20)-C(29)	110(1)
C(1)-C(10)-C(9)	104 (1)	C(21)-C(20)-C(30)	114 (1)
C(1)-C(10)-C(25)	107 (1)	C(29)-C(20)-C(30)	109 (1)
C(5)-C(10)-C(9)	106 (1)	C(20)-C(21)-C(22)	112 (1)
C(5)-C(10)-C(25)	116 (1)	C(20)-C(21)-O(11) C(22)-C(21)-O(11)	108-5 (9)
C(9)-C(10)-C(25)	115 (1)	C(22)-C(21)-O(11)	106-4 (9)
C(9)-C(11)-C(12)	114 (1)	C(17)-C(22)-C(21)	117 (1)
C(11)-C(12)=C(13)	125 (1)	C(17)-C(22)-O(9)	107-2 (9)
C(12)=C(13)-C(14)	121 (1)	C(21)-C(22)-O(9)	105-4 (9)
C(12)=C(13)=C(14)	121 (1)	C(21) C(22) C(3)	,
(ii) Extranualear			
(ii) Extranuclear	122 (1)	O(4)-C(33), C(34)	121 (2)
C(24)-O(3)-C(24)	123 (1)	O(4)=C(33)-C(34)	131 (2)
C(16)-O(5)-C(35)	121 (1)	O(5)-C(35)=O(6)	120 (1)
C(28)O(7)C(28)	119 (1)	O(5)-C(35)-C(36)	111 (1)
C(21)-O(11)-C(1')	116-3 (9)	O(6)=C(35)-C(36)	129 (2)
C(4)-C(24)-O(3)	109 (1)	O(7)-C(37)=O(8)	116 (2)
C(17)-C(28)-O(7)	109 (1)	O(7)-C(37)-C(38)	113 (2)
O(1)-C(31)=O(2)	125 (2)	O(8)=C(37)-C(38)	129 (2)
O(1)-C(31)-C(32)	106 (2)	O(9)-C(39)=O(10)	125 (2)
O(2)=C(31)=C(32)	128 (2)	O(9)-C(39)-C(40)	104 (1)
O(3)-C(33)=O(4)	115 (2)	O(10)=C(39)-C(40)	130 (2)
O(3)-C(33)-C(34)	114 (2)	0(10)=0(0)) 0(10)	(-,
0(3) 0(33) 0(34)	114 (2)		
(iii) Chaonasanosal moi	at 11		
(iii) Glucopyranosyl moi		C(2')O(4')C(12')	117(1)
O(11)-C(1')-C(2')	110 (1)		
O(11)-C(1')-O(1')	108.8 (9)	O(2')-C(7')-O(3')	119 (2)
C(2')-C(1')-O(1')	109 (1)	O(2')-C(7')-C(8')	109 (1)
C(1')-C(2')-C(3')	110 (1)	O(3')-C(7')-C(8')	132 (2)
C(1')-C(2')-C(4')	110(1)	O(2")C(7")O(3")	126 (2)
C(3')-C(2')-O(4')	106-3 (9)	O(2")-C(7")-C(8")	110(1)
C(2')-C(3')-C(4')	110(1)	O(3")-C(7")-C(8")	124 (2)
C(2')-C(3')-O(2'')	104 (1)	C(7')-C(8')=C(9')	112 (2)
C(4')-C(3')-O(2')	108 (1)	C(7')-C(8')-C(11')	118 (2)
C(4')-C(3')-O(2') C(3')-C(4')-C(5')	112 (1)	C(9')=C(8')-C(11')	130 (2)
C(3')-C(4')-C(3') C(3')-C(4')-O(2')	108 (1)	C(7'')-C(8'')=C(9'')	129 (2)
		C(7")-C(8")-C(11")	106 (2)
O(2')-C(4')-C(5')	104 (1)		125 (2)
C(4')-C(5')-C(6')	113 (1)	C(9'')=C(8'')-C(11'')	
C(4')-C(5')-O(1')	108 (1)	C(8')=C(9')-C(10')	134 (2)
C(8')-C(5')-O(1')	108 (1)	C(8'')=C(9'')-C(10'')	132 (2)
C(1')-O(1')-C(1')	111-6 (9)	O(4')-C(12')=O(5')	125 (1)
C(4')-O(2')-C(7')	120 (1)	O(4')C(12')C(13')	112(1)
C(3')-O(2'')-C(7'')	114 (1)	O(5')=C(12')-C(13')	123 (2)

In the triterpene skeleton, several C-C bonds [C(1)-C(10); C(5)-C(10); C(8)-C(14); C(9)-C(10);C(17)-C(22) deviate from the mean value of 1.54 Å. it seems significant that all these long C-C bonds are attached to at least one of the fully substituted carbon atoms [C(8), C(10), C(14) and C(17)]. Several C-C-C angles also deviate from the accepted value of 109.3°. The degrees of substitution of the carbon

atoms and long-range steric strain in the structure (s below) seem to be responsible for these deviations.

The C(12)–C(13) bond [1.37 (2) Å] is shorter that the remainder of the ring-system bonds and com sponds to the expected location of the double bond oleanene triterpenes.

In Table 4, the conformation of the ring systemi analyzed in terms of the least-squares planes an interplanar angles. The equations of the least-square planes were calculated by the method of Schomako @(6)-(Waser, Marsh & Bergman (1959). The torsional angle (5)-(in the ring system are given in Table 5(i). From Table (6)—(4 and 5, it is clear that the rings A, B, D and E take

Table 4. Least-squares planes

C(14)-**C**(9)-(**E**(11)–

chair confc **C**(12 rings E rin distor distor action C(10 Other the re

(a) Equations of the planes
Plane A C(1), C(2), C(3), C(4), C(5), C(10)
-0.8682x - 0.4203y - 0.2637z + 9.6570 = 0
Plane B $C(5)$, $C(6)$, $C(7)$, $C(8)$, $C(9)$, $C(10)$
-0.8343x - 0.5346y + 0.1344z + 9.2884 = 0
Plane C C(8), C(9), C(11), C(12), C(13), C(14)
-0.7409x - 0.6612y + 0.1178z + 7.8474 = 0
Plane D C(13), C(14), C(15), C(16), C(17), C(18)
-0.8455x - 0.5273y - 0.0838z + 10.3836 = 0
Plane E C(17), C(18), C(19), C(20), C(21), C(22)
-0.8856x + 0.1556y + 0.4376z + 9.1570 = 0
Plane $F \ C(21), O(11), C(1')$
0.0951x - 0.9953y - 0.0203z - 2.6655 = 0
Plane G C(1'), C(2'), C(3'), C(4'), C(5'), O(1')
-0.4992x - 0.6567y + 0.5653z + 1.0736 = 0

	0 1,7,23			0.00	ber	tw(
(b) Dev	iations (Å) from the le	ast-squares plar	nes	val	
•	•				dis	
Plane A		Plane	В	Plane C		
C(1)	-0.24 (1) C(5)	-0.50(1)	C(8)	0.38 (I) me	
C(2)	0.25 (2	2) C(6)	0.15(1)	C(9)	_0.29 (I) dia	
C(3)	-0.24 (2	2) C(7)	-0·15 (1)	C(11)	0.04 (2) the	re
C(4)	0.22 (1) C(8)	0.39(1)	C(12)	0.09(1)	·C
C(5)	-0 ⋅23 (1	1) C(9)	-0.08(1)	C(13)	-0.004 (Th
C(10)	0.23 (1			C(14)	-0.23(4	
O(1)	0⋅26 (1	l) C(25)	1.66 (2)	C(26)	1.88 (1) CO	
C(23)	-0.56 (2	2) C(26)	1.89 (1)	C(27)	-1.77 (l) (at	C
C(24)	1.69 (1				int	er:
C(25)	1.75 (2	2)			sho	111/
					. Cin	
Plane L)	Plane		Plane G		
C(13)	0.11 (1		-0·13 (1)	C(1')	_0.26 (I) tov	va;
C(14)	-0.12(1	l) C(18)	0.18(1)	C(2')	0·22 () C(28
C(15)	0.17 (1	l) C(19)	-0.27(1)	C(3')	-0.21 (le rin	29
C(16)	-0.23 (1	l) C(20)	0.30(1)	C(4')	U-21 (e .
C(17)	0.21 (1	C(21)	0·26 (1)	C(5')	-U·24 (·//	
C(18)	-0.16(1		0.19(1)	O(1')	0.28 (i) fin	
C(27)	-1.63(1	, ,	0.83(1)	O(11)	0·15(axi	a]
O(5)	-1.71(1	C(29)	0·29 (1)	O(4')	-0·40 (!) C(22
C(28)	1.75 (1	(30) C(30)	1.81 (1)	O(2")	0.54 (J. mu	ele
		O(11)	0.28(1)	O(2')	-U-32 (Day	in.
		· O(9)	-0.52 (1)	C(6')		
						TŁ
(c) Inter	planar an				Çoı	
	A/B	156.0 (4)	D/E	129.	1 (3) 🔑 Çqi	ıa [.]
	B/C	170-9 (4)	E/F		6 (2) Wal	lie.
	C/D	164-9 (4)	F/G	126-	5 (3)	u (

Table 5. Torsional angles (°)

repene nucleus		C(8)-C(9)-C(11)-C(12)	37.0(1)	(ii) Glucopyranosyl ring	
C(1)-C(2)-C(3)	55.6 (2)	C(9)-C(11)-C(12)-C(13)	6.5 (1)	O(1')-C(1')-C(2')-C(3')	58-6 (2)
G(2)-C(3)-C(4)	58-1 (2)	C(11)-C(12)-C(13)-C(14)	2.0(1)	C(2')-C(1')-O(1')-C(5')	$-67 \cdot 1 (3)$
C(3)-C(4)-C(5)	54.8 (2)	C(12)-C(13)-C(14)-C(15)	-150.0(3)	C(1')-C(2')-C(3')-C(4')	-54.2 (2)
(4)-C(5)-C(10)	-53-6 (2)	C(18)-C(13)-C(14)-C(15)	32.3 (2)	C(2')-C(3')-C(4')-C(5')	-53·0 (2)
C(5)-C(6)-C(7)	64.9 (2)	C(14)-C(13)-C(18)-C(17)	−35·8 (2)	C(3')-C(4')-C(5')-O(1')	-56.9(2)
(5)-C(10)-C(9)	-165.5(4)	C(13)-C(14)-C(15)-C(16)	37.4 (2)	C(4')-C(5')-O(1')-C(1')	63.8 (3)
G(5)-C(10)-C(1)	171.9 (3)	C(14)-C(15)-C(16)-C(17)	48-5 (3)		
C(5)-C(10)-C(9)	61.1 (2)	C(15)-C(16)-C(17)-C(18)	50-4 (3)	(iii) Angelyl groups	112 4 (2)
C(6)-C(7)-C(8)	60.3 (2)	C(16)-C(17)-C(18)-C(13)	42.9 (3)	C(4')-C(3')-O(2'')-C(7'')	113.4 (3)
C(1)-C(8)-C(9)	50.4 (2)	C(16)-C(17)-C(18)-C(19)	83.5 (3)	O(2")-C(7")-C(8")-C(9")	-11.60(2)
C(8)-C(9)-C(10)	50.5 (2)	C(22)-C(17)-C(18)-C(13)	$-169 \cdot 1 (3)$	O(3")-C(7")-C(8")-C(11")	-19.54(2)
7°C(8)-C(9)-C(11)	179 · 7 (4)	C(22)-C(17)-C(18)-C(19)	42.6 (3)	C(11")-C(8")-C(9")-C(10")	171·14 (3) 106·8 (4)
M-C(8)-C(9)-C(10)	169-1 (3)	C(18)-C(17)-C(22)-C(21)	43.1 (2)	C(3')-C(4')-O(2')-C(7')	144.3 (4)
M-C(8)-C(9)-C(11)	-61.7(2)	C(17)-C(18)-C(19)-C(20)	56.7 (3)	O(2')-C(7')-C(8')-C(9')	144.3 (4)
I-C(8)-C(14)-C(13)	56.3 (1)	C(18)-C(19)-C(20)-C(21)	-62.2(3)	O(3')-C(7')-C(8')-C(11')	174.6 (4)
II)-C(9)-C(10)-C(1)	61.5 (2)	C(19)-C(20)-C(21)-C(22)	59.6 (3)	C(11')-C(8')-C(9')-C(10')	17470 (4)

tr'conformation while the C ring adopts a half-chair formation as a result of the double bond between (12) and C(13). The junctions of the A/B and B/C \blacksquare are in the trans form while that between the D and mage is cis. The rings B, D and E are significantly lotted as is the whole molecular skeleton. These fortions seem to be caused mainly by the intertions between the bulky axial groups (at C(4), C(8), (0) on one hand and at C(14) and C(16) on the which push each other away in order to release repulsive forces. Actually, the non-bonded distances ween 1,3 diaxial groups are larger than the expected in non-distorted rings. We have reported these Inces in Table 6. In a regular chair-form sixmbered ring, the non-bonded distances between 1,3 mill groups would be 2.52 A. These distances are inbered ring, the non-bonded distances between 1,3 increased to, for example, 3.27 (2) Å [C(24) C(25)].

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C

0.38(1)

-0.29 (1)

0·09 (1) -0.004 (1)

-0·23 (1) 1·88 (1)

-0.26 (1)

0·22 (1) -0·21 (1)

0·21 (1) -0·24 (1) 0·28 (1)

0.15(1)

0.54 (I) -0.52 (I)

0.28 (2)

!9·1 (3)

15.6 (2)

The distortion of ring C from a regular half-chair formation may be due to the degree of substitution C(8) and C(14). Also resulting from 1,3 diaxial action is the distortion of the whole molecule. As no by the angles between the mean planes of the A, B, C and D, the triterpene skeleton is convex and the axial groups C(24), C(25), C(26) and D. On the other hand there is twisting between E and D about the long direction of the molecule. The same previous of the three bulky acetate groups situated at C(16), C(17) and A similar conformation of the triterpenic was has been previously observed in gymnematory Hoge & Nordman (1974).

glucopyranosyl ring has an almost regular chair of the substituent groups in lorial positions. The bond distances agree with the reported for β -D-glucose (Chu & Jeffrey, 1968)

Table 6. Some non-bonded distances (Å) between 1,3 diaxial groups

C(24)C(25)	3.27(2)
C(25)C(26)	3.20(2)
C(27)O(5)	3.06(2)
O(5)O(9)	2.97 (1)

Table 7. Shortest intermolecular distances (Å) with standard deviations in parentheses

Key to symmetry operations relating designated atoms to reference atoms at (x,y,z):

y, ž	(iv) $x + 1, \frac{1}{2} + y, z + 1$ (v) $x, y - 1, z$ (vi) $\bar{x} + 2, \frac{1}{2} + y, \bar{z} + 2$.		
$y, \hat{z} + 1$			
3.56 (3)	C(30)···O(2 ¹¹¹)	3.63 (2)	
3.69(2)	$C(32)\cdots O(4^{iv})$	3.10(3)	
3.60(2)	C(34)···O(2 ⁿ)	3.30(3)	
3.35(2)	$C(36)\cdots C(40^{tl})$	3.57 (3)	
3.60(2)	$C(38)\cdots O(4^{l})$	3.58 (3)	
3.56(2)	$O(1)\cdots O(8^l)$	3.36 (2)	
	$O(6)\cdots C(3^{\prime li})$	3.46 (2)	
	$O(6)\cdots C(5'^{li})$	3.68 (2)	
3.55 (2)	$C(13')\cdots O(3'^{vl})$	3.54 (3)	
	3.56 (3) 3.69 (2) 3.60 (2) 3.35 (2) 3.56 (2) 3.56 (2) 3.52 (3) 3.61 (3)	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	

with one exception: the C(3')–C(4') bond is significantly shorter in comparison with the accepted value of 1.54 Å. Moreover, the two angles C(2')–C(3')–O(2'') and C(5')–C(4')–O(2') differ from the tetrahedral angle of 109.5° . Steric strain resulting from the two long angelyl side chains situated at C(3') and C(4') could be responsible for these unusual values. Indeed, these angelyl groups, in order to overcome any overlapping, are rotated with respect to each other by

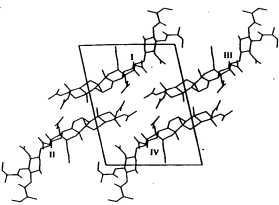


Fig. 3. Projection of the acetylated napoleogenin structure along the b axis. The positions of the molecules are; (I) at x, y, z; (II) at $\bar{x} + 1$, $\frac{1}{2} + y$, z; (III) at x, y, z + 1; (IV) at x + 1, $\frac{1}{2} + y$, $\bar{z} + 1$ with x, y and z coordinates given in Table 1.

about 180° about the C(7')—C(8') bond [the corresponding torsional angles are reported in Table 5(iii)].

One should also notice that in both angelyl and acetyl groups, the accuracy of bond lengths and valency angles is highly restricted by a large thermal motion of the end-chain atoms.

A packing diagram of the crystal structure viewed along the b axis is shown in Fig. 3. The intermolecular distances less than 3.7 Å are reported in Table 7. There is no abnormal feature in the packing, which appears to

be dictated mainly by close van der Waals contained between the molecules.

The authors wish to thank Professors J. Touss amin and A. Van de Vorst for their interest in this work in dependent of Mr M. Vermeire for technical assistance. Crystals acetylated napoleogenin were provided by the H Laboratoire de Chimie Organique of Professor R. H.

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The Structure of 2-Amino-5-nitrophenol: a Comparison with 2-Aminophenol

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Abstract

The crystal structure of 2-amino-5-nitrophenol has been determined from visually estimated Cu $K\alpha$ data, and refined to an R value of 0.072 for 1138 non-zero reflexions. The crystals are monoclinic, $P2_1/c$, with a=8.49 (1), b=10.34 (1), c=8.05 (1) Å, $\beta=113\cdot1$ (1)° and Z=4. The molecules, related by a glide plane, are held together by $O-H\cdots NH_2$ hydrogen bonds $[O\cdots N 2.801$ (3), $H\cdots N 1.89$ (4) Å, $O-H\cdots N 174$ (3)°] to form a ribbon along c. The ribbons are held together by dipole—dipole interactions

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between the molecules related by I, as found monoly in mononitrophenols. The morphotropism mononitrophenols and aminophenols is discussed.

Introduction

As part of a programme of studies on the crystal molecular structures of phenols (Kagawa, Kashino & Haisa, 1976) and the systemization organic crystals (Haisa, 1978), the structure amino-5-nitrophenol has been determined in order

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